

FILE 'USPAT' ENTERED AT 12:55:14 ON 23 JUL 1999

*Excerpt from full
transcript including
each sketchy*

6. 5,648,247, Jul. 15, 1997, Method for increasing the omega-hydroxylase activity in candida tropicalis; Stephen Picataggio, et al., 435/142, 254.22 [IMAGE AVAILABLE]
8. 5,620,878, Apr. 15, 1997, Method for increasing the omega-hydroxylase activity in Candida tropicalis; Stephen Picataggio, et al., 435/142, 254.22 [IMAGE AVAILABLE]
L1 0 S PICHIA PASTORIS (P) CYTOCHROME P450 (P) (TRANSFORM?
OR T
RAN
L2 0 S PICHIA PASTORIS (P) MONOOXYGENASE (P) (TRANSFORM? OR
TRA
NSF
L3 9 S (YEAST OR CANDIDA MALTOSA) (P) CYTOCHROME P450 (P)
(TRAN
SFO
L4 47145 S ALKANE HYDROXYLAT? OR DICARBOXYL?
L5 13 S L4 AND CYTOCHROME P450
L6 11 S L5 NOT L3
L7 3 S POX4 AND URA3
L8 70 S CANDIDA MALTOSA
L9 48 S L8 AND HOST CELL
L10 39 S L9 AND HETEROLOG?
L11 1 S CANDIDA MALTOSA /TI

U.S. Patent & Trademark Office LOGOFF AT 13:13:53 ON 23 JUL 1999

FILE 'HOME' ENTERED AT 13:13:32 ON 23 JUL 1999

=> file medline, biosis, caplus, agricola

L1 ANSWER 1 OF 2 MEDLINE
AN 96154241 MEDLINE
DN 96154241
TI Functional expression of recombinant spiny dogfish shark (Squalus acanthias) cytochrome P450c17 (17 alpha-hydroxylase/C17,20-lyase) in yeast (Pichia pastoris).
AU Trant J M
CS Department of Zoology and Physiology, Louisiana State University, Baton Rouge 70803, USA.. trant@umbi.umd.edu
SO ARCHIVES OF BIOCHEMISTRY AND BIOPHYSICS, (1996 Feb 1) 326 (1) 8-14.
Journal code: 6SK. ISSN: 0003-9861.
CY United States
DT Journal; Article; (JOURNAL ARTICLE)
LA English
FS Priority Journals; Cancer Journals
EM 199605
AB The cDNA encoding the spiny dogfish shark (Squalus acanthias) testicular

form of cytochrome P450c17 (CYP17) was used to direct the
 heterologous
 expression of a functional enzyme in yeast (***Pichia***
 pastoris). This protein possesses two enzymatic
 activities: 17
 alpha-hydroxylase and C17,20-lyase reactions. Cytochrome P450c17
 is a key
 steroidogenic enzyme for the production of sex steroids in gonadal
 tissue
 and for cortisol production in adrenal tissue. This study
 describes the
 culture conditions and the enzymatic activity of
 recombinant
 shark cytochrome P450c17. The shark enzyme was compatible with the
 endogenous yeast NADPH- ***cytochrome*** ***P450***
 reductase and
 was bioactive within the living yeast cell. Progesterone (at 15
 microM)
 was metabolized (51 pmol/min/10(9) cells) faster than pregnenolone
 (36
 pmol/min/10(9) cells). Both progesterone and pregnenolone were
 completely
 metabolized to their respective androgens (androstenedione and
 dehydroepiandrosterone). Although 11 beta-hydroxy-progesterone was
 readily
 17 alpha-hydroxylated by the shark P450, the lyase reaction was
 not
 evident. Alterations to the 2-carbon sidechain of progesterone
 (21-hydroxylation or 20 beta-reduction) prevented metabolism.
 High-density
 cultures (> 1.5 x 10(9) cells/ml) yielded the greatest quantity of
 recombinant protein but cultures of lower density
 produced more
 recombinant protein per cell. This is the first report
 of
 heterologous expression in yeast of a steroidogenic
 cytochrome
 P450 from a lower vertebrate.

L6 ANSWER 9 OF 13 CAPLUS COPYRIGHT 1999 ACS

AN 1988:524518 CAPLUS

DN 109:124518

TI Degradation of long-chain n-alkanes by the yeast ***Candida***
 maltosa . II. Oxidation on n-alkanes and intermediates

using

microsomal membrane fractions

AU Blasig, R.; Mauersberger, S.; Riege, P.; Schunck, W. H.; Jockisch,
 W.;

Franke, P.; Mueller, H. G.

CS Cent. Inst. Mol. Biol., Ger. Acad. Sci., Berlin, DDR-1115, Ger.
 Dem. Rep.

SO Appl. Microbiol. Biotechnol. (1988), 28(6), 589-97
 CODEN: AMBIDG; ISSN: 0175-7598

DT Journal

LA English

AB Microsomal membrane fractions of the yeast C. maltosa were
 investigated

with respect to their ability to catalyze the oxidn. of n-alkanes, fatty alcs. and fatty acids. Anal. of intermediates of n-hexadecane oxidn. led to the conclusion that monoterminal attack was predominant, whereas diterminal oxidn. proceeded as a minor reaction. The oxidn. of long-chain primary alcs. to the corresponding aldehydes occurred without addn. of NAD (phosphate) [NAD(P)+] and was accompanied by stoichiometric oxygen consumption and hydrogen peroxide prodn., suggesting that an alc. oxidase instead of an NAD(P)+-requiring alc. dehydrogenase catalyzed these reactions. As shown for n-hexadecane, the hydroxylation of palmitic acid was found to be carbon monoxide-dependent, indicating involvement of a cytochrome P 450 system, as in the case of n- ***alkane*** ***hydroxylation***.

L6 ANSWER 10 OF 13 CAPLUS COPYRIGHT 1999 ACS

AN 1987:210770 CAPLUS

DN 106:210770

TI Function and regulation of cytochrome P-450 in alkane-assimilating yeast.

II. Effect of oxygen-limitation

AU Schunck, W. H.; Mauersberger, S.; Kaergel, E.; Huth, J.; Mueller, H. G.

CS Cent. Inst. Mol. Biol., Ger. Acad. Sci., Berlin-Buch, DDR-1115, Ger. Dem. Rep.

SO Arch. Microbiol. (1987), 147(3), 245-8
CODEN: AMICCW; ISSN: 0302-8933

DT Journal

LA English

AB Transition of n-hexadecane utilizing cultures of ***Candida*** ***maltosa*** to oxygen-limited growth caused an .ltoreq.6-fold increase

of the cellular cytochrome P 450 content. Enhanced cytochrome P 450

formation required protein de novo synthesis and was not due to a change

of the apo/holo-enzyme ratio as demonstrated by cycloheximide inhibition

and immunol. quantitation. The effect of low oxygen concn. (pO₂ = 3-5%)

was simulated by selective inhibition of ***alkane***

hydroxylation with carbon monoxide (at a pO₂ of 70-75%).

Enhanced

cytochrome P 450 formation occurred even when a const. growth rate was

maintained through utilization of a second nonrepressive growth substrate.

However, the presence of n-alkanes was an essential precondition.

Apparently, the cytochrome P 450 formation was mainly regulated by the

intracellular inducer concn. which depends on the relative rates
of alkane
transport into the cell and the actual ***alkane***
hydroxylating activity of the enzyme system.
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(FILE 'HOME' ENTERED AT 13:13:32 ON 23 JUL 1999)

FILE 'MEDLINE, BIOSIS, CAPLUS, AGRICOLA' ENTERED AT 13:14:54 ON 23
JUL 1999

L1 2 S PICHIA PASTORIS (P) CYTOCHROME P450 (P) (TRANSFORM? OR
TANSFEC
L2 0 S PICHIA PASTORIS (P) MONOOXYGENASE (P) (TRANSFORM? OR
TRANSFEC
L3 55430 S ALKANE HYDROXYLAT? OR DICARBOXY?
L4 75 S L3 AND CYTOCHROME P450
L5 13 S L4 AND CANDIDA MALTOSA
L6 13 S L5 NOT L1
L7 5 S POX4 AND URA3
L8 5 S L7 NOT L1
L9 5 S L7 NOT L6

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